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## AMR-meta: ak-mer and metafeature approach toclassify antimicrobial resistance fromhigh-throughput short-read metagenomics data --Manuscript Draft--

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Abstract:	genes through comparison with curated current methods are often hampered by homology/homoplasy with other non-AN Results. We present AMR-meta, a data on k -mers, which combines algebraic m regularized regression. Metafeatures ca antibiotic classes. In addition, AMR-meta joins an AMR gene database with non-Acompare AMR-meta with AMRPlusPlus, their ensemble via a voting system. In co (interquartile) f-score of 0.7 (0.2-0.9). Or test on average AMR-meta yields a 1.3 In terms of run-time, AMR-meta is 3x fast MARC, and as fast as AMRPlusPlus. Fin ontologies and observed variance of all development on standardization of bench Conclusions. AMR-meta is a fast, accurate sets to improve sensitivity and specificity	microbial samples enables profiling of AMR AMR databases. However, performance of database incompleteness, and presence of MR genes in sequenced samples.  Abase-free and alignment-free approach, based natrix factorization into metafeatures with puture multi-level gene diversity across main a employs an augmented training strategy that AMR genes (used as negative examples). We a DeepARG, and Meta-MARC, further testing ross-validation, AMR-meta has a median an semi-synthetic metagenomic dataexternal 3-fold hit rate increase over existing methods. Seter than DeepARG and 30x faster than Metanally, we note that differences in AMR tools in classification outputs call for further chmarking data and protocols.  The differences in AMR ontologies and the noutputs call for the deployment of standard				
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PAPER

# AMR-meta: a *k*-mer and metafeature approach to classify antimicrobial resistance from high-throughput short-read metagenomics data

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#### **Abstract**

**Background** Antimicrobial resistance (AMR) is a global health concern. High-throughput metagenomic sequencing of microbial samples enables profiling of AMR genes through comparison with curated AMR databases. However, performance of current methods are often hampered by database incompleteness, and presence of homology/homoplasy with other non-AMR genes in sequenced samples.

Results We present AMR-meta, a database-free and alignment-free approach, based on *k*-mers, which combines algebraic matrix factorization into metafeatures with regularized regression. Metafeatures capture multi-level gene diversity across main antibiotic classes. In addition, AMR-meta employs an augmented training strategy that joins an AMR gene database with non-AMR genes (used as negative examples). We compare AMR-meta with AMRPlusPlus, DeepARG, and Meta-MARC, further testing their ensemble via a voting system. In cross-validation, AMR-meta has a median (interquartile) f-score of 0.7 (0.2-0.9). On semi-synthetic metagenomic data -external test- on average AMR-meta yields a 1.3-fold hit rate increase over existing methods. In terms of run-time, AMR-meta is 3x faster than DeepARG and 30x faster than Meta-MARC, and as fast as AMRPlusPlus. Finally, we note that differences in AMR ontologies and observed variance of all tools in classification outputs call for further development on standardization of benchmarking data and protocols.

**Conclusions** AMR-meta is a fast, accurate classifier that exploits non-AMR negative sets to improve sensitivity and specificity. The differences in AMR ontologies and the high variance of all tools in classification outputs call for the deployment of standard benchmarking data and protocols, to fairly compare AMR prediction tools.

Key words: functional metagenomics; short reads; antimicrobial resistance; machine learning; matrix factorization

#### Introduction

Antimicrobial resistance (AMR) is the ability of microorganisms to resist the effect of drugs targeted to eliminate them [1], and is globally recognized as a threat to public health as it

makes treatment of microbial infections harder, increasing the risk of disease spread and severity [2]. Data from 890 U.S. hospitals collected on specific combinations of antibiotics and bacteria show that AMR caused an estimated 622,390 infections in 2017 [3]. Treating infections caused by AMR is clinically chal-

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#### **Key Points**

- · AMR-meta is a novel, database-free and alignment-free approach, combining matrix factorization with a training strategy including an AMR gene database plus non-AMR genes.
- On cross-validated results, AMR-meta has a median f-score of 0.7, while on external test sets it yields a 1.3-fold hit rate increase over existing methods.
- AMR-met is 3x to 30x faster than state-of-the art algorithms.

lenging since it requires to identify which drugs the infecting strain is susceptible to, and then to take a timely decision on the therapy to use. Notably, AMR is not limited to healthcare, as it represents a significant challenge also in animal and plant health, and thus in the entire ecosystem [4]. Therefore, detecting AMR in clinical, veterinarian, and botanical isolates is pivotal to curb the spread of AMR pathogens and reduce its impact. Although culture-based methods can accurately detect AMR, they are resource intensive with respect to trained personnel, monetary cost, and time [5]. Moreover, since only a fraction of bacterial species are cultivable with standard methods, culturebased methods are only applicable to a small number of bacteria. For these reasons, whole genome and metagenomics sequencing has become an increasingly prevalent method for AMR characterization. The challenge that then arises is how to accurately identify and quantify the AMR genes from such sequencing data. To accomplish this, a number of different methods have been proposed. Despite the concordance between in silico genotypic and in vitro phenotypic resistance assessment, the uptake of AMR prediction tools for routine healthcare has been slow, and they showed discordant performance in clinical settings [6].

AMR prediction methods for metagenomics rely on comparison to databases of AMR genes. Two comprehensive and widely used of AMR databases are the Comprehensive Antibiotic Resistance Database (CARD) [7, 8] and MEGARes [9, 10]. CARD is thoroughly maintained, with monthly updates on AMR determinants that have (i) an associated peer-reviewed scientific publication, (ii) a DNA sequence available in GenBank, (iii) clear experimental evidence of elevated minimum inhibitory concentration over controls. Currently, CARD integrates over 3,000 reference sequences of AMR genes and over 1,500 single nucleotide polymorphisms, knowledge on resistance mechanisms, and specific antibiotic classes. CARD uses a manually curated process and ontology, named the Antibiotic Resistance Ontology (ARO, github.com/arpcard/aro), which describes the molecular relations of antibiotic resistance (e.g., acquired resistance genes, drug targets, AMR mechanisms). MEGARes [9] -and its most recent 2.0 update [10]- is a hand-curated AMR database designed for high throughput sequencing data processing. MEGARes includes CARD genes and variants, but utilizes a different annotation structure. Specifically, it is a multi-level hierarchy (type, mechanism, class, group) in the form of a direct acyclic graph, ensuring that two higher level ranks are not linked to the same lower level rank. MEGARes annotation graph is therefore an optimal structure for ecological profiling and construction of AMR classifiers because, for example, it cannot result in conflicting sequence classification. MEGARes 2.0 currently includes ~8,000 genes. Major improvements from its first release consist in the inclusion of antibacterial biocide and metal resistance genes.

For AMR classification of metagenomic samples from highthroughput sequencing, one class of methods is based on the use of sequence read aligners. One widely used tool in this category is AMRPlusPlus [9], which aligns all reads to MEGARes using Burrows-Wheeler Aligner (BWA) [11] and then postprocesses the alignment to identify the genes that have over 80% coverage from the alignment, providing the associated AMR annotation in the output. AMRPlusPlus 2.0 [10] is an improved version of AMRPlusPlus that is designed to be faster for large-scale projects. AMRPlusPlus 2.0 provides a postalignment classification through the ResistomeAnalyzer (quality measure for nucleotide coverage of a reference sequence for a given read) and the RarefactionAnalysis (assessment of sequencing depth) modules. It also incorporates prediction of AMR due to single nucleotide polymorphisms in housekeeping genes, using a curated set that matches CARD. Of note, CARD also performs AMR prediction for housekeeping genes via the Resistant Gene Identifier (RGI), available as a web-service and a command-line application. Although alignment-based methods have high precision [12], they can only classify reads which align to known AMR genes. Given that existing AMR databases are incomplete, a large portion of novel AMR genes may go undetected.

Another class of methods for AMR characterization is alignment-free, employing a variety of approaches including substring (k-mer) matching and machine learning. ResFinder [13] and KmerResistance [14] process metagenomic reads by first constructing the set of all unique k-length subsequences (called k-mer spectrum) from the dataset. ResFinder 4.0 compares the set of unique k-mers to detect AMR genes and AMRrelated chromosomal gene mutations based on an reference database built on a collection of chromosomal point mutations in bacterial pathogens [15], resistance genes from the Antibiotic Resistance Genes Database (ARDB) [16] and other literature sources [17]. The user is required to input a specific bacterial species for which the resistance is searched. Eight bacterial species are available. KmerResistance, as ResFinder, compares the set of unique k-mers to an ad hoc gene AMR reference database derived from literature [18, 19]. Specifically, Kmer-Resistance uses exact co-occurring k-mer matching between a query sequence and the database, with a "winner takes all" strategy, i.e., multiple k-mer occurrences on different genes are resolved by selecting the one with highest frequency. Next, a quality measure of a whole AMR gene match is defined as a probability function of coverage (i.e., fraction of the genome covered by at least one k-mer) and depth (i.e., average number of times the k-mers in the match). Similar to alignmentbased methods, ResFinder and KmerResistance are also bound to identifying genes that are found in a specified database, and therefore, have limited ability to detect putative AMR sequences. Another limitation of the *k*-mer based approaches is the low flexibility with respect to sequencing errors [14], possibly increasing false negative rates in sequence classification.

Other alignment-free methods use machine learning classifiers to identify putative and known AMR genes, such as Resfams [20] and Meta-MARC [12], both based on hierarchical hidden Markov models (HMMs). Resfams [20] preprocesses high-throughput sequences by assembling them and translating the resulting contigs into amino acid sequences. Meta-MARC can predict AMR for an input sequence (either a short read or a longer assembled contig), according to the resis-

tance class, group, and mechanism hierarchy defined in the MEGARes hierarchical data structure. Specifically, Meta-MARC is an ensemble of HMMs, each trained on a group of genes from MEGARes. A classification is performed by aggregating predictions from the lowest level of the MEGARes annotation hierarchy towards the highest level. Meta-MARC achieves better sensitivity, specificity, fraction of classified high-throughput sequence data, and number of AMR classes identified when compared to alignment matches and Resfams. However, the performance of Meta-MARC with short read data is worse than classifying assembled contigs.

DeepARG [21] is a hybrid machine learning and alignmentbased approach that leverages convolutional deep learning networks. The alignment module first translates the input sequences to amino acids and using DIAMOND [22], and then aligns the translated sequences to a custom AMR database created by merging CARD, ARDB [16], and manually selected AMR sequences from the Universal Protein Resource (UNIPROT). The deep learning model then predicts the AMR class for all aligned reads. Since the machine learning step is subsequent to the alignment one, de facto DeepARG suffers from the limitations of alignment-based AMR prediction algorithms.

For completeness, it is worth mentioning AMR gene identification methods that are not specifically designed for highthroughput short read metagenomic data. These methods take as input one or a combination of: single genes, specific genome strains, genomic or proteomic variants, and/or protein primary, secondary, or tertiary structures. Similar to the methods described previously, these methods use alignment and/or machine learning paradigms [23, 24, 25, 26, 27, 28, 29, 30]. These algorithms bind the user into performing one or more supplementary pre-processing steps on metagenomics data, not included into the algorithm, such as sequence alignment or assembly, sequence translations into proteins, or protein structure prediction. Because of the required pre-processing, these methods defy the very advantages provided by the alignmentfree design. For further reference, Hendriksen et al. [31] provide a comprehensive review.

In this paper, we develop AMR-meta, a novel, alignmentfree, AMR classification approach for high-throughput metagenomic data, based on k-mers and matrix factorization of kmers. The matrix factorization produces a number of 'metafeatures' able to capture multiple levels of gene diversity within broad AMR classes. Importantly, and differently from existing methods, AMR-meta uses an augmented training strategy that incorporates non-AMR genes as negative examples. We show that our approach is competitive with state-of-the-art tools (i.e., AMRPlusPlus 2.0, Meta-MARC, and DeepARG) in classification performance and execution speed. Notably, AMR-meta captures resistance mechanics complementary to those found by other tools, which instead are more correlated to each other.

#### **Methods**

AMR-meta is trained and tested first on an internal dataset that -differently from other approaches- includes both AMR (named resistant) and non-AMR genes (named susceptible). The AMR genes are taken from MEGARes 2.0 [10], while non-AMR genes are chosen from Genbank's RefSeq and include (a) bacterial genes that are highly dissimilar to AMR genes, and (b) AMR-homologous sequences, i.e. sequences highly similar to AMR genes, but not known to be associated to antibiotic resistance. By including the non-AMR and AMR-homologous sequences, we aim to decrease the false positive calls and to increase the true negative rates. This internal dataset is split into a 70/30 training/test ratio, and AMR-meta components (k-mers and k-mer-derived metafeatures) are trained

and tested accordingly (all performance measures reported in this paper are relative to test sets). Second, we generate two semi-synthetic external datasets, drawing bacterial genomes from the Pathosystems Resource Integration Center (PATRIC) [32], and simulating short read data. We derive two PATRIC datasets that represent drug resistance/susceptibility relative to specific molecules or antibiotic classes, called  ${\mbox{PSS}}_{mol}$  and PSS<sub>cla</sub>, respectively. This twofold design allows us to benchmark AMR-meta against other existing tools -AMRPlusPlus 2.0, Meta-MARC, and DeepARG- in a a flexible way, since their outputs levels vary among antibiotic classes and more specific mechanisms. We use  $\ensuremath{\mathsf{PSS}_{mol}}$  to score the AMR predictions, and PSS<sub>cla</sub> to estimate the concordance of AMR-meta class predictions with those of other methods. Finally, we combine AMRmeta with the other tools, and evaluate their predictions on two functional metagenomic datasets that were sampled a clinical and environmental setting. Our internal/external workflow is summarized in Figure 1.

#### Feature encoding and prediction models

#### AMR-meta k-mer LASSO module

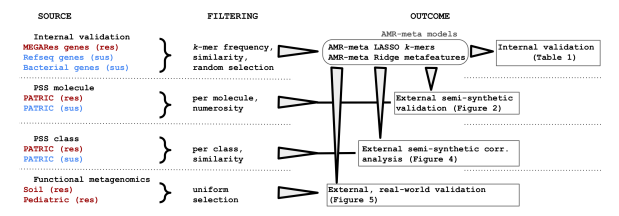
The baseline models of AMR-meta are logistic regressors one for each antibiotic class– that use raw k-mers as input. Each model utilizes the whole class-specific *k*-mer spectrum (derived from the collated positive/negative training datasets), where each feature is a binary value, representing the presence or absence of a particular *k*-mer in the dataset. Given the high-dimensionality of the k-mer spectrum, we use least absolute shrinkage and selection operator (LASSO) regularization to reduce the feature space, optimizing the shrinkage operator via cross-validation [33]. Given the heterogeneity in gene diversity within each class, e.g. Betalactamases have higher diversity than Floroquinolones, we also expect different cardinality of non-zero coefficients among the class-specific k-mer LASSO regressors.

#### AMR-meta metafeature ridge module

One possible problem with k-mer LASSO regression is that a single linear combination of k-mer features might not be able to explain the variance of the entire dataset, even if discrimination performance is good for the majority of genes in one class. A way to increase the portion of variance explained is to use more than one linear combination, e.g., the first m-th vectors of a principal component analysis. In this way, multiple independent combinations of k-mers can more effectively represent the genetic diversity within antibiotic classes.

Accordingly, we explore a space transformation -with concomitant dimension reduction- of the k-mer spectrum that identifies a set of (orthogonal) multiple features, i.e., metafeatures, each as an independent combination of the original kmers contributing to a cumulative portion of the data variance. To do so, we apply a matrix factorization approach, which has been previously shown apt to tackle complex feature extraction problems, e.g., oncology and proteomics [34, 35]. The method is based on non-negative matrix tri-factorization [36]. The algorithm identifies low-rank, non-negative matrices whose product provides an approximation of the original non-negative matrix.

Here we consider two data domains, namely k-mers and genes. A k-mer is related to a gene if it is present in the gene sequence. Let us denote the total number of genes with g; the total number of k-mers with t; a matrix of rrows and c columns having all values equal to zero with  $\varnothing_{r,c}$ ; and a matrix with one gene per row, and one k-mer per column  $R_{g,t}$  with, and  $R_{g,t}^T$  as its transpose. We denote the transpose of a matrix A with superscript T as  $A^T$  in the



**Figure 1.** AMR-meta training/test workflow. We assemble an internal dataset of AMR and non-AMR homologous genes from MEGARes and RefSeq genes, on which AMR-meta models (k-mers, and metafeatures through matrix factorization) are trained and tested (70/30 split). AMR-meta and other AMR classification tools are then externally tested on: (i) semi-synthetic data from PATRIC at both antibiotic class and molecule levels (PSS<sub>cla</sub> and PSS<sub>mol</sub>); and (ii) functional metagenomics data (Soil and Pediatric).

rest of this paper. We express the relation between the two domains by a symmetrical, four-block matrix  $R = \binom{\varnothing g,g}{R_{g,t}^T} \binom{R_{g,t}^T}{N_{g,t}}$ , where non-diagonal block matrices represent the relation (intersections) between k-mers and genes. Note that in this context, the relation between elements is defined by design: We set the value of a R at an entry to 1 if the corresponding k-mer is present in the corresponding gene, and 0 otherwise.

We denote the number of k-mer metafeatures and the number of gene metafeatures as  $m_t$  and  $m_g$ , respectively. The factorization procedure decomposes R into the product of three matrices G, S, and  $G^T$ , such that  $G \times S \times G^T$  will approximate R by reducing the error up to a user-defined lower bound set as the difference between two consecutive iterations (denoted with  $R \approx GSG^T$ ). Here G represents the relation between the original domains (genes, k-mers) and their metafeatures; and S represents the relation between the metafeatures, i.e., how one domain is mapped to the other. The matrices G and S have the following form both expressed as four block matrices:

$$\mathsf{G} = \left( \begin{smallmatrix} G_{g,m_g} & \varnothing_{g,m_t} \\ \varnothing_{k,m_g} & G_{t,m_t} \end{smallmatrix} \right) \text{ and } \mathsf{S} = \left( \begin{smallmatrix} \mathsf{S}_{m_g,m_g} & \mathsf{S}_{m_g,m_t} \\ \mathsf{S}_{m_t,m_g} & \mathsf{S}_{m_t,m_t} \end{smallmatrix} \right).$$

We use the intersection between the data of the same domain as constraints in the factorization process, i.e., each domain has a block, symmetrical constraint. We define the matrix  $\Theta$  to represent the self-domain relations, i.e., gene/gene and k-mer/k-mer relations. Therefore,  $\Theta$  is an  $R\times R$  matrix. The empty blocks of  $\Theta$  are the non-diagonal blocks.  $\Theta=({{\Theta g,g}\atop{\mathscr{O}_{t,g}}}{{\mathscr{O}_{t,t}}\atop{\mathscr{O}_{t,g}}}).$ 

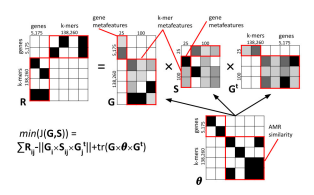
In  $\Theta$  we set each entry to -1 if the corresponding row and column elements share a relation; 1 if unrelated; and 0 if the relation is unknown. In this application, in the  $\Theta_t$  block we consider each k-mer identical to itself (related, -1), while we make no assumption about the relation with two different k-mers (not related, 0). In the  $\Theta_g$  block, we consider all the genes of each class to be related (-1), and different from the genes of other classes (1).

The goal of the factorization is to minimize the following objective function:

$$min_{G\geq 0}(G;S) = \sum ||R_{ij} - G_i S_{ij} G_j^t|| + tr(G \Theta G^t)$$
 (1)

where  $||\cdot||$  indicates the Frobenius norm, and  $tr(\cdot)$  indicates the trace. The objective function is composed of two parts: The first part measures the difference between the original matrix and the product of the three factorized matrices; the second

part calculates the adherence of the factorized metafeatures to the constraints, in our case based on the AMR resistance class. The factorization process proceeds in an iterative fashion until convergence to a local minimum, with convergence heuristically defined by observing the value of the objective function and the corresponding reconstruction error below a user-defined threshold [36, 34, 35]. We fix a threshold of  $10^{-2}$ as the difference between consecutive iterations, or reaching 5,000 iterations, as stop criteria. Previous works discuss the method in detail [34, 35]; a dedicated github repository contains code and user manual github.com/smarini/MaDDA. The factorization process, calculated over the full length training genes, produces  $G_{t,m_t}$ , which is the matrix relating the k-mers to their metafeatures. For each short read pair encoded as binary vector of k-mer occurrences  $sr_{1,t}$ , we calculate its metafeatures as  $sr_{1,t} \times G_{t,m_t}$ . Since the optimal number of metafeatures can be hard to infer, and the sizes of the matrices grow with the number of features [34, 35], for this application we used up to  $m_t$  = 100 and  $m_q$  = 25 metafeatures. After factorization, we feed the metafeatures to a logistic regression, optimizing the coefficients with a ridge approach. Figure 2 provides a graphical representation of the factorization process.



**Figure 2.** The matrix tri-factorization scheme. AMR, non-AMR, and AMR-homologous genes are paired up with k-mers across all antibiotic resistance classes into the R matrix, and the the dimension is reduced through the  $R \approx GSG^t$  factorization, where the metafeatures are extracted, revealing the AMR similarity phenotypes in the  $\theta$  matrix.

#### **Training strategy**

#### AMR genes.

We collate AMR genes from MEGARes 2.0 [10], constituting the positive (resistant) reference sets on the basis of the MEGARes annotation at the antibiotic class level. Of note, we exclude housekeeping genes that confer resistance through single point mutations.

#### Putative non-AMR bacterial genes.

We include putative non-AMR genes from the RefSeq database [37]. Using BLAST, we select the 1,000 RefSeq bacterial genes that do not match to MEGARes (e-value=10), aiming for a 1:1 target ratio with the antibiotic class of highest frequency. This gene set has high genetic divergence from the AMR genes in MEGARes, yet the nucleotide content is fully bacterial.

#### AMR-homologous human and vertebrate genes.

To mimic genes that likely do not provide AMR, but share a significant similarity with AMR genes we assemble a dataset selecting AMR-homologous genes and gene fragments from the human genome (GRCh38), and all the contigs in RefSeq labelled as 'vertebrate mammalian' and 'vertebrate other' assemblies. To do so, we run an ungapped BLAST search of all MEGARes genes against these human and vertebrate sequences (e-value=0.01). We use each unique sequence match, and add the flanking region to each match, elongating the matched sequence to be equal in length to the corresponding resistant MEGARes gene. Specifically, with a match of  $n_{match}$  nucleotides between target and query AMR gene, we extend the match by  $\frac{n_{match}}{2}$  nucleotides in both directions on the target MEGARes sequence. The underlying assumption here is that matches of bacterial AMR genes on vertebrate genomes are spurious or not functional, and therefore do not provide AMR. Of note, this setup is similar to the test set derivation presented in DeepARG

#### k-mer based and metafeature modelling.

All k-mers present in the genes of the training datasets, excluding any sample reserved for validation (see next subsection), are considered and counted using different values of k, from 13 to 77 based on prior literature evidence [14]. The best value for *k* is chosen incrementally on the basis of internal validation performance, stopping when performance decreases. Next, we stratify the training samples by class. We remove all *k*-mers with a frequency less than a given cut-off f in a single class (3 or 5 upon internal validation). We also exclude AMR classes with with less than 10 k-mers after frequency filtering.

#### Simulation of metagenomic short read data for training.

We use the AMR datasets described above to generate short reads, labelling each as resistant or susceptible to an antibiotic class. For each MEGARes class, we generate short read datasets providing 10x base coverage of the original full-gene data. These datasets allow the evaluation of both false positives and false negatives.

#### **External validation**

We use four independent external datasets, two semi-synthetic (made similarly to the training set), and two from functional metagenomic experiments. As the prevalence of AMR and the k-mer spectrum in the external test set is not guaranteed to be balanced as in the training, we re-calibrate the k-mer and metafeature probability threshold for external validation using the internal validation dataset and a number of samples where the k-mer and metafeature vectors are empty, i.e., they represent the non-AMR gene background. The ratio is optimized between 1:0.05 and 1:10, picking the first that meets the calibration target, i.e., a prediction with a score < 0.5 for a feature vector without any *k*-mer belonging to our model.

#### Semi-synthetic datasets.

We create the semi-synthetic datasets from PATRIC, downloading via FTP full bacterial genomes and summary metadata [30, 23]. We retain only genomes annotated as susceptible or resistant after an antibiogram test conform to the Clinical & Laboratory Standards Institute (CLSI), which is the most frequent testing standard in PATRIC, with over 55,000 resistant and 54,000 susceptible records [30]. Since the antibiotic nomenclature in PATRIC is molecule-specific and does not match exactly the MEGARes ontology hierarchy, we compile a lookup table linking each PATRIC drug annotation to a MEGARes class. We remove PATRIC genomes that do not refer to the AMR classes considered in the training phases, or are not included in the classes predicted by the concurrent methods.

We then generate two PATRIC semi-synthetic datasets (PSS), based on PATRIC antibiotic molecule labels (PSS<sub>mol</sub>) and MEGARes classes (PSS $_{cla}$ ), respectively.

We use  $PSS_{mol}$  to assess the performance of our approach and the concurrent methods on molecule-specific data. We retain genomes are resistant (or susceptible) to at least one MEGARes class. We rank the PATRIC drug labels based on number of associated genomes, and we select the top ones based on the associated MEGARes classes. We exclude labels with less than 250 genomes, or labels not referring to a specific molecule (e.g., Tetracycline). We generate 250,000 short reads for each PATRIC label, equally divided between resistant and susceptible. Note that for  ${\ensuremath{\mathsf{PSS}}}_{mol},$  as the PATRIC labels refer to genome (and not the specific gene, as in MEGARes), it is not possible to determine the ground truth, i.e., if a short read comes from a resistant or a susceptible gene. To assess methods' performances, in absence of such ground truth, we develop a scoring system based on the assumption that a method should find more resistant read pairs from resistant genomes, and less from susceptible genomes. With sr<sub>res,res</sub> defined as the number of short read pairs coming from resistant genomes and classified as resistant, and with sr<sub>res,sus</sub> as the number of short read pairs coming from susceptible genomes and classified as resistant, we define the S-score as  $S = sr_{res,res} - sr_{res,sus}$ . A higher S-score thus denotes better performance, and a negative value implies that the method finds more resistant short read pairs among the susceptible ones.

 $\mathrm{PSS}_{cla}$  is collated at the class level. Unlike  $\mathrm{PSS}_{mol}$ , each short read from  ${\mbox{PSS}}_{cla}$  has a known label which indicates if it comes from a resistant or susceptible gene. To generate PSS<sub>cla</sub>, first we remove PATRIC genomes presenting inconsistent class annotations, i.e., that are annotated as both resistant and susceptible to antibiotics belonging to the same class. Second, in order to consider only genomes that are resistant (or susceptible) to the range of antibiotics within a given MEGARes class, we rank each genome in decreasing order of the total number of annotations of resistance (or susceptibility) to multiple drugs within the same class. Based on this ranking, we retain only genomes that rank over the 90th percentile. Third, we perform a class-by-class BLAST filtering (e-value=0.01, percent identity ∈ [70,90]) of the selected PATRIC genomes against MEGARes genes, retaining and clipping the unique genes of PATRIC genomes that match MEGARes. The objective is to extract a set of PATRIC genes that match to MEGARes genes, but are not exact matches. In fact, genes similar to known resistant genes coming from antibiotic susceptible -by a phenotypic test- genomes represent excellent candidates to test the ability of classifier to recognize true/false positives. From these selected PATRIC genes, we generate short reads covering the

selected genes, and capping the number of resistant or susceptible paired reads up to 100,000 per AMR class (i.e., 400,000 total reads per class). We reckon that with this procedure, we are able to label uniquely each PATRIC instance that passes the filter; however, in the BLAST alignment, there could be flanking regions or inserts that produce artifact matches. Nonetheless, given the strict parameters used, we we deem these cases to be rare. A resistant sample likely contains only resistant reads, and vice-versa for a susceptible sample. Therefore, it is possible to calculate sample-wide performance by counting the proportion of resistant-within-resistant and susceptiblewithin-susceptible reads in each test sample. After filtering, Glycopeptides and Lipopeptides are excluded as there are less than fifteen resistant genomes. Sulfonamides are excluded as no susceptible genomes is retained by our filtering procedure.

#### Functional metagenomics data.

We benchmarked our method against two functional metagenomic datasets, which we refer to as the Pediatric and the Soil datasets (NCBI BioProject Accessions PRJNA244044 and PR-JNA215106). A functional metagenomics experiments is made by cloning metagenomic DNA fragments into bacterial vectors grown on antibiotic-laden media. The cultured bacteria surviving the antibiotic are sequenced using a clonally amplified highthroughput sequence library. As per experimental design, for each fosmid, all sequence reads contain at least one AMR gene (known or not yet discovered) resistant to a known antibiotic. Therefore, each sequencing experiment has a known antibiotic resistance label. However, since the original metagenomics fragments can be longer than a single AMR gene, a single fosmid might contain multiple AMR genes, or contain unknown genes. The Pediatric and Soil datasets include fosmids from Escherichia coli (DH10B) and consist of of 219 and 169 samples with an average of 1.98 and 1.12 million paired-end short reads respectively, sequenced with Illumina Genome Analyzer IIx technology. We utilize the aforementioned PATRIC annotation lookup table to pair antibiotic annotations to MEGARes classes. For testing classifiers' performance, we randomly select 100,000 short read pairs for each class as for the PATRIC datasets.

#### Software and hardware setup

We process the training/validation data, the semi-synthetic  $\ensuremath{\mathsf{PSS}_{mol}}$  and  $\ensuremath{\mathsf{PSS}_{cla}}$  datasets, and the experimental functional metagenomics data through in house UNIX scripts, off-theshelf bioinformatics tools including BLAST, R (r-project.org/), and Bioconductor (bioconductor.org/). The k-mer LASSO and the metafeature regression are developed in R, bash, and C++. We download the functional metagenomics datasets using NCBI's sra-toolkit. For short read generation, we use InSilicoSeq [38], simulating Illumina's NovaSeq (company's top-line production scale sequencing instrument) reads with default parameters. We exclude genes shorter than 151 bases (length of NovaSeq's short reads) from the simulations. Code and R scripts are available publicly at github.com/smarini/AMR-meta under the MIT license.

#### **Results**

#### AMR-meta provides competitive prediction performance on multiple AMR classes

We generate thirteen datasets, corresponding to the antibiotic classes (according to the MEGARes ontology) of: Aminoglycosides, Betalactamases, Drug and biocide resistance,

Fluoroquinolones, Glycopeptides, Lipopeptides, Macrolide-Lincosamide-Streptogramin (MLS), Multi-biocide resistance, Multi-drug resistance, Multi-metal-resistance, Phenicols, Sulfonamides, and Tetracyclines. We exclude classes with less than 10 k-mers after frequency filtering. Upon internal validation, the best k-mer length k and frequency threshold f are 13 and 5, respectively (the performance decreases at k=31 and for f=3 with the same or higher k). Upon optimization of the kvalue, the total number of unique 13-mers is 138,260, and the median (interquartile range, IQR) number per class is 3,645 (1,658-7,168). The matrix factorization includes 5,175 training genes, yielding a matrix R of 138,260 + 5,175 = 143,435 rows and columns, and a k-mer/metafeature matrix of 138, 260  $\times$  100 elements.

Table 1 shows the class-specific performance summaries by k-mer and metafeature regression on the internal validation sets. On the internal validation set, the k-mer LASSO and the metafeature regression exhibit a good tradeoff between sensitivity and specificity at both k values. The median (IQR) number of features selected by k-mer LASSO is 12,783 (12,304 and 13,179). As expected, the highest number of non-zero coefficients is found in the Betalactamase class, which is the class with higher diversity and number of resistant genes in MEGARes. The same holds for the highest number of metafeatures with positive coefficients (note that each metafeature is derived from the matrix factorization described above, incorporating several hundred thousands *k*-mer/gene elements). In terms of performance, for LASSO, the median (IQR) f-measure across all classes is 0.7 (0.2-0.9), while for the metafeature regression, the median f-measure is 0.4 (0.2-0.7). For both methods, the best performing classes are Betalactamases, and Fluoroquinolones, while the most problematic are MLS, and Multi-biocide, -drug, and -metal resistance. Despite the kmer LASSO having a higher median f-measure, the metafeature regression performs better in the problematic MLS and Drug and biocide classes, shows better sensitivity in Glycopeptides, and better Specificity in Fluoroquinolones and Lipopetides. For reference comparison, the median (IQR) f-measure across classes is 0.5 (0.3-0.7) for DeepARG, and 0.9 (0.9-1.0) for Meta-MARC, based on the original papers' validation results. AMRPlusPlus 2.0 does not report per-class results on test sets.

#### AMR-meta generalizes robustly on external, semisynthetic datasets

The  $PSS_{mol}$  dataset includes twelve molecule labels incorporated into antibiotic classes, namely: ciprofloxacin and levofloxacin (Fluoroquinolones); gentamicin and amikacin (Aminoglycosides); ceftriaxone and ampicillin (Betalactamases); chloramphenicol (Phenicols); sulfisoxazole (Sulfonamides); erythromycin and azithromycin (MLS); tigecycline (Tetracyclines); and vancomycin (Glycopeptides). Performance results in terms of S-score, which summarizes the correct resistance and susceptible hits (the higher the better), are shown in Figure 3. The median (IQR) S-score for the k-mer LASSO is 285.5 (123.5, 540), and for the metafeature regression is 322 (73, 470). Meta-MARC scores 250 (72, 359.5), DeepARG scores 144.5 (43, 345), and AMRPlusPlus 2.0 scores -29 (-377.5, 210). Overall, our metafeature approach shows both the highest performance and stability, exhibiting also a positive score in the levofloxacine molecule, whereas all the other methods produce a negative score. The k-mer LASSO component ranks second, followed by the other off-the-shelf tools.

Table 1. Performance of k-mer LASSO and metafeature ridge regression in predicting antibiotic class susceptibility/resistance on the internal test sets (30% of full dataset). Results show f-measure, Matthew's correlations coefficient (MCC), sensitivity and specificity; also, the number of non-zero k-mer LASSO and positive metafeature ridge coefficients are shown.

		k-mer LASSO			Metafeature ridge						
Antibiotic Class	N (test)	#feat.	F-measure	MCC	Sens.	Spec.	#metaf.	F-measure	MCC	Sens.	Spec.
Aminoglycosides	4,920	13,162	0.85	0.84	0.79	0.99	54	0.58	0.54	0.57	0.97
Betalactamases	36,052	19,483	0.96	0.93	0.94	0.99	74	0.89	0.79	0.83	0.96
Drug and biocide resistance	5,055	13,064	0.36	0.39	0.93	0.76	56	0.39	0.93	0.7	0.66
Fluoroquinolones	1,286	11,462	0.98	0.98	0.96	1	50	0.9	0.9	0.92	1
Glycopeptides	3,200	12,700	0.8	0.8	0.7	1	54	0.23	0.27	0.84	0.75
Lipopeptides	1,084	12,356	0.85	0.85	0.76	1	43	0.8	0.8	0.73	1
Macrolide-Lincosamide-Streptogramin	2,210	14,064	0.2	0.28	0.93	0.77	54	0.3	0.29	0.38	0.97
Multi-biocide resistance	1,412	12,304	0.13	0.2	0.88	0.76	51	0.1	0.16	0.78	0.73
Multi-drug resistance	1,387	12,280	0.13	0.21	0.91	0.77	48	0.11	0.18	0.83	0.74
Multi-metal resistance	2,407	13,179	0.21	0.28	0.92	0.76	62	0.18	0.25	0.9	0.73
Phenicols	922	11,115	0.74	0.74	0.66	1	51	0.44	0.44	0.53	0.99
Sulfonamides	531	12,783	0.75	0.78	0.6	1	54	0.75	0.77	1	0.6
Tetracyclines	4,208	14,286	0.86	0.85	0.8	1	43	0.67	0.65	0.67	0.98

#### AMR-meta predictions complement those of existing algorithms

Next, we measure the correlation between the predictions of the two AMR-meta modules and the ones from the other algorithms. As  $PSS_{mol}$  does not have a per-gene defined ground truth, we assemble  $PSS_{cla}$ . The  $PSS_{cla}$  dataset includes six out of the thirteen MEGARes classes, namely Aminoglycosides, Betalactamases, Fluoroquinolones, MLS, Phenicols, and Tetracyclines. PSS<sub>cla</sub> has instances from both positive (resistant) and negative (susceptible) genes. When we look at the classspecific concordance for each pair of tools using the the Spearman's rank correlation (Figure 4),  ${
m PSS}_{cla}$  shows that the algorithms behave differently. Specifically, DeepARG, Meta-MARC, and AMRPlusPlus 2.0 are highly correlated in most of antibiotic classes (range 0.59-0.92), while they have low correlation with the k-mer LASSO and the metafeature regression (range 0.04-0.12) -which in turn show mild-low correlation (range 0.12-0.49). Thus, both k-mer LASSO and metafeature regression stand distant from each other and the other methods. The PSS<sub>cla</sub> dataset is explicitly constructed to measure classspecific concordance, with very similar resistant and susceptible instances. However, for this reason, the  $PSS_{cla}$  becomes by design a challenging dataset for classification, because the reads derived from susceptible genes all well align with other resistant genes in the same AMR class. Thus, the performance

of all algorithms will tend to flatten. Nonetheless, the metafeature approach exhibits the highest median accuracy. Overall -pooling both resistant and susceptible for each AMR classthe k-mer LASSO median (IQR) percent of correct predictions is 44% (35%-48%), the metafeature ridge 46% (33%-48%), DeepARG 44% (36%-47%), AMRPlusPlus 2.0 45% (36%-50%), and Meta-MARC 44% (36%-47%).

#### AMR-meta ensemble for functional genomics

The Soil and Pediatric datasets come from functional metagenomics experiments that by design guarantee the presence of antibiotic resistance in a sequence sample, since the sample is cultured on antibiotic-laden medium. However, sequenced reads can also contain other or unknown genes, which cannot be quantified. We consider here the hit rate, i.e. the proportion of sequence reads classified as resistant. Cautionary, a higher hit rate can signify that a method finds more AMR genes, but also that a method finds more false positives. Given that AMRmeta is designed to decrease false positive rate, we expect it to be the most conservative. Yet, in order to identify empirically a tradeoff between the approaches, in addition to running each single model, we also built an ensemble using voting with k-mer LASSO, the metafeature regression, and the individual models' predictions as input features (requiring at least two concordant predictions for classifying resistance).

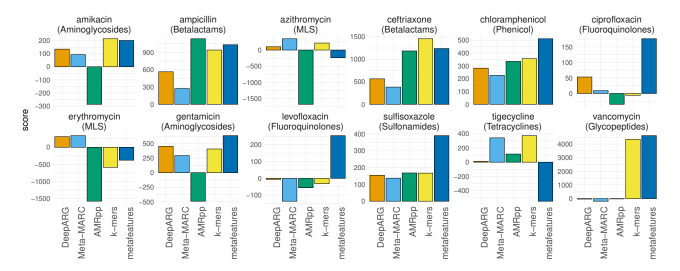
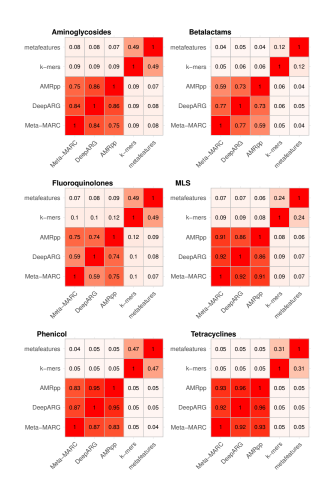


Figure 3. Performance of AMR-meta (k-mers and metafeatures) and of other off-the-shelf tools on the molecule-level PATRIC semi-synthetic data (PSS $_{mol}$ ). The S-score score is the difference between short read pairs predicted as resistant from the pooled resistant and susceptible genomes drawn from PATRIC.



**Figure 4.** Spearman's Rank correlation of the AMR classifiers on the PATRIC semi-synthetic data ( $PSS_{cla}$ ).

On Soil, the voting ensemble achieves the highest hit rate with a median (IQR) fraction of read pairs identified as resistant of 7.72% (1.28%-10%), followed by AMRPlusPlus 2.0 with 7.03% (1.06%-7.48%), DeepARG with 6.27% (1.21%-7.32%), Meta-MARC with 4.97% (1.86%-8.68%), the k-mer approach with 1.94% (0.7%-2.49%), and the metafeature approach with 0.08% (0.01%-0.65%). On Pediatric, Meta-MARC achieves the highest hit rate with a median (IQR) of 8.51% (2.29%-28.14%), followed by the k-mer approach with 0.27% (0.2%-4.8%), the voting ensemble with 0.27% (0.05%-4.97%), AMRPlusPlus 2.0 with 0.2% (0.02%-11.95%), DeepARG with 0.19% (0.02%-8.06%), and the metafeature approach with 0.01% (0%-0.4%). We observe large variations in each method depending on the class considered. It has to be noted that Meta-MARC's threshold was previously re-calibrated on these datasets, and its standard threshold is much more conservative. As expected, the metafeature module is the most conservative on both datasets, while the voting ensemble offers a balanced alternative in all cases. Interestingly, the k-mer approach is one of the least conservative on the Pediatric set. Detailed results on the external Pediatric and Soil functional metagenomics datasets are illustrated in Figure 5.

#### Run-time comparison

To compare execution times, we create benchmark datasets of increasing size by selecting reads drawn the semi-synthetic PATRIC data (across all classes), generating files of 1GB, 2GB, 5GB, and 10GB of paired short read files. We run all algorithms on University of Florida's High Performance Cluster –

HiPerGator 3.0– using a single node, composed by four AMD Opteron 6378 cores, with 32GB of RAM. Table 2 show run times on the node. AMRPlusPlus 2.0 and MetAMR *k*-mer LASSO are the fastest tools, with a time of execution difference within minutes up to 5GB load. DeepARG is up to 3 times slower than MetAMR, and Meta-MARC is considerably slower (30–folds), hitting the 24–hour wall time for files larger than 1GB.

#### **Discussion**

In this work, we present AMR-meta, an alignment-free, *k*-mer- and metafeature-based AMR classifier for short read metagenomics data. AMR-meta uses an augmented training strategy based on non-AMR and AMR-homologous genes, providing relevant classification performance increment across various antibiotic classes.

Historically, the main objective of AMR characterization by metagenomics sequencing has been to identify known AMR genes, using comprehensive and up-to-date databases. However, the absence of non-AMR genes (negative examples) and of AMR-homologous sequences that do not have role in resistance can hamper AMR classification accuracy, and impact the trade-off between sensitivity and specificity. Notably, there are metagenomics classification tools that exploited the negative-positive *k*-mer representation paradigm. For instance, Clark weighs differently *k*-mers that are found only in specific species, as compared to those that are shared by different species or genuses [39]. Other studies, focused on full-genome analysis and based on in vitro susceptibility, have shown high discriminating ability and capacity to identify potential new resistance features [27, 29].

It is worth to mention that comparing different AMR tools can be challenging, because not all use the same ontology, or provide classifications at the same annotation level. For instance, Meta-MARC is trained on a self-determined similaritybased clustering of AMR genes, yet it is able to provide predictions at the mechanism/class/group level according to MEGARes ontology, matching the outputs of AMRPlusPlus 2.0 and AMR-meta. Instead, DeepARG uses a unique set of AMR categories derived from the CARD and ARDB. At this point, comparison of tools requires making an arbitrary choice on the AMR ontology to be used, and on the annotation level (e.g. class rather than mechanism), potentially penalizing one approach over another, as we show in our semi-synthetic PATRIC datasets PSS<sub>mol</sub> and PSS<sub>cla</sub>. In addition, summarizing results over antibiotic classes can also introduce bias, given the high class imbalance in terms of antibiotics, gene frequency, and the aforementioned heterogeneity of intra-class gene diversity. It is understandable that a unified AMR ontology is difficult to achieve, yet an effort of the community to create common, standardized protocols for benchmarking and comparison is

One limitation of our approach is in the sample resistance/susceptibility annotation for validation and benchmark datasets. First, we label most of bacterial genes that do not match to MEGARes as drug-susceptible, whilst in reality these sequences might contain new, undiscovered AMR genes. Second, there might be inconsistencies with antibiogram results in PATRIC.

Other limitations include the fact that we try only one metafeature approach –matrix factorization– while other methods could be tested, e.g., sparse binary principal/independent component analysis. Finally, it is known that k-mer approaches are not very sensitive to mutations, while mutant genes can still carry resistance.

Future development for AMR-meta includes new strategies to select positive/negative labelled examples (and mu-

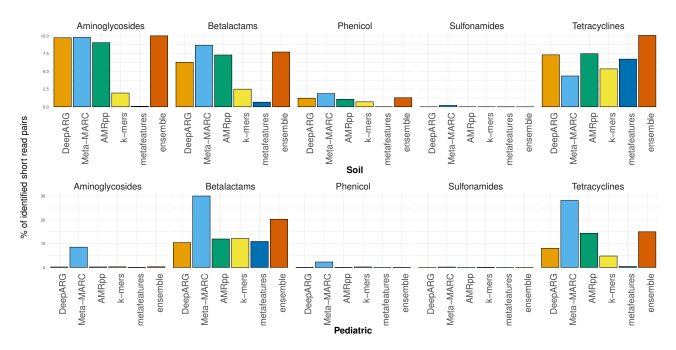


Figure 5. Percentage of sequence reads predicted resistant on the functional metagenomics data (Soil and Pediatric) by AMR-meta, off-the-shelf tools, and their voting ensemble.

Table 2. Running times (hh:mm:ss) of AMR classification tools on metagenomics short read data (reads drawn from the PATRIC datasets), 151 bases, paired end, fastq format).

File size (R1+R2)	# of reads (R1+R2)	AMR-meta (k-mer)	AMRPlusPlus 2.0	Meta-MARC	DeepARG
1GB	1,584,451	00:22:16	00:21:19	16:26:27	00:53:01
2GB	3,168,014	00:43:37	00:49:40	>24h	01:38:55
5GB	7,924,402	01:47:24	01:35:47	>24h	03:41:06
10GB	15,851,366	03:32:46	02:48:43	>24h	11:42:16

tant genes) can further improve the classification performance. As another perspective, given the availability of efficient data structures for k-mer modelling, the LASSO module of AMRmeta could also be efficiently implemented as standalone AMR classifier to process data from portable sequencers in real time using mobile devices [40].

#### Availability of source code and requirements

· Project name: AMR-meta

• Project home page: https://github.com/smarini/AMR-meta

Operating system(s): Linux

Programming language: Bash, R, C++

Other requirements: R packages Matrix, stringr, glmnet

· License: MIT

#### Availability of supporting data and materials

As stated in the Methods, the data sets supporting the results of this article are obtainable from public, ncbi.nlm.nih.gov/refseq; specifically Refseq, MEGARes, megares.meglab.org; NCBI BioProject PRJNA244044, ncbi.nlm.nih.gov/bioproject/244044; NCBI BioProject PRJNA215106 ncbi.nlm.nih.gov/bioproject/215106; and PATRIC, patricbrc.org. AMR-meta algorithm, including a containerized version via Singularity, is available at github.com/smarini/AMR-meta.

#### List of abbreviations

AMR - antimicrobial resistance ARDB - Antibiotic Resistance Genes Database BWA - Burrows-Wheeler Aligner CARD - Comprehensive Antibiotic Resistance Database DIAMOND - double index alignment of next-generation sequencing data LASSO - least absolute shrinkage and selection operator PATRIC -Pathosystems Resource Integration Center IQR - interquartile range HMM - hidden Markov model UNIPROT - Universal Protein Resource

#### **Ethical Approval**

Not applicable.

#### Consent for publication

Not applicable

#### **Competing Interests**

The authors declare that they have no competing interests.

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#### **Author's Contributions**

SM, MP, CB, NRN, and TK conceived the idea and wrote the paper. SM, MO, IS, and RAD prepared the data and performed the experiments. All authors read and approved the final manuscript.

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Dear Editorial Office,

we are pleased to submit "AMR-meta: a k-mer and metafeature approach to classify antimicrobial resistance from high-throughput short-read metagenomics data" to Gigascience.

Please note that this is a resubmission—this manuscript was initially submitted with the ID GIGA-D-21-00255. The manuscript was returned with the invitation to provide a containerized version and resubmit:

We appreciate the work you have presented ... If you're able to containerize this work so it is easily deployable and testable, we would be willing to reconsider this work for GigaScience. Please update the manuscript with the relevant information and resubmit an updated version.

Following the recommendations on this communication, we implemed a containerized version and resubmitted. We updated both the github and the manuscript, which now reads "AMR-meta, including a containerized version via Singularity, is available at <a href="https://github.com/smarini/AMR-meta">https://github.com/smarini/AMR-meta</a>" in the "Availability of supporting data and materials" section.

The singularity container is hosted by the ICBR institute (University of Florida) servers. ICBR committed to maintain the download link available for at least five years.

AMR-meta is a novel --database-free and alignment-free-- approach to classify antimicrobial resistance (AMR) from high-throughput metagenomics data. Using algebraic matrix factorization, our approach infers multiple 'metafeatures' able to capture gene diversity across antibiotic classes with multiple resistance mechanisms. Importantly, AMR-meta employs an augmented training strategy that joins an AMR gene database with putative non-AMR and AMR-homologous genes (used as negative examples). Thus, unlike alignment-based methods, AMR-meta is not limited to identifying only known AMR genes.

We compared AMR-meta with other popular AMR classification tools --AMRPlusPlus, DeepARG and MetaMARC-- showing its competitive advantage in terms of performance and, especially, runtime, making it suitable for large-scale processing.

We believe that AMR-meta is a great fit for Gigascience, and especially for the Functional Metagenomics collection. We look forward to receiving the Editor's feedback.

In case reviewers are needed, we recommend the following experts:

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#### Sincerely,

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